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AMENDMENTS TO THE CLAIMS

This listing of claims will replace all prior versions and listings of claims in the

application:

LISTING OF CLAIMS:

1. (Currently amended) A method for isolating and culturing multipotent

progenitor/stem cells from cord blood-derived isolated mononuclear cells, which comprises

culturing the cord blood-derived isolated mononuclear cells successively in:

1) a first animal cell culture medium comprising fetal bovine serum_(FBS), L-

glutamine and granulocyte macrophage-colony stimulating factor_(GM-CSF), in addition to

inorganic salts, vitamins, and amino acids and/or supplementary elements;

a second animal cell culture medium which is the same as the first animal cell

culture medium except for lacking GM-CSF; and

3) a third animal cell culture medium which is the same as the first animal cell

culture medium except that GM-CSF is replaced with stem cell factor $\underline{\hspace{0.1cm}}(SCF)$ and endothelial

growth factor_(EGF).

2. (Currently amended) The method of claim 1, wherein the animal cell culture

medium further contains D-glucose ranging from 3,500 to 5,500 $\underline{mg/\ell}$ $\underline{mg/M\ell}$ and sodium

pyruvate ranging from 50 to 200 mg/l mg/Ml.

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(Currently amended) The method of claim 1, wherein the first animal cell culture

medium contains 10 to 20% FBS, 1 to 2 mM L-glutamine, \underline{and} 10 to 100 ng/ \underline{ml} HE GM-CSF; the

second animal cell culture medium contains 10 to 20% FBS and 1 to 2 mM L-glutamine; and the

third animal cell culture medium contains 10 to 20% FBS, 1 to 2 mM L-glutamine, 10 to 100

ng/ml mg/Mℓ-SCF and 10 to 50 ng/ml mg/Mℓ EGF.

(currently amended): The method of claim 1, wherein the cultivation in the first

animal cell culture medium is conducted by inoculating the mononuclear cells into the first

animal cell culture medium at a concentration of 1×10⁵ to 1×10⁶ cells/cm² and culturing at 37°C

under an atmosphere of 5% CO2 for 1 to 2 weeks; the cultivation in the second animal cell

culture medium is conducted by replacing the first animal cell culture medium by the second

animal cell culture medium after confirming the formation of a multi-layer cell colony and

further culturing at 37°C under an atmosphere of 5% CO₂ for 1 to 2 weeks; and the cultivation in

the third animal cell culture medium is conducted by inoculating the cells cultured in the second

animal cell culture medium into the third animal cell culture medium at a concentration of 2×10⁴

 ${\color{red}to~8\times10^4~cells/cm^2} - {\color{red}1\times10^5~to~4\times10^5~cells/ml}~after~observing~the~metamorphosis~of~the~multi-layer$

cell colony into a mono-layer cell colony and further culturing at 37°C under an atmosphere of

5% CO2 for 1 to 2 weeks.

5. (Currently amended) A multipotent progenitor/stem cell isolated and cultured

from a cord blood-derived isolated mononuclear cell, the multipotent progenitor/stem cell being

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is isolated and cultured by a method comprising culturing the cord blood-derived mononuclear cells successively in:

- a first animal cell culture medium comprising fetal bovine serum (FBS), Lglutamine and granulocyte macrophage-colony stimulating factor (GM-CSF), in addition to inorganic salts, vitamins, and amino acids-and/or supplementary elements;
- a second animal cell culture medium which is the same as the first animal cell culture medium except for lacking GM-CSF; and
- 3) a third animal cell culture medium which is the same as the first animal cell culture medium except that GM-CSF is replaced with stem cell factor_(SCF) and endothelial growth factor_(EGF),

wherein said multipotent progenitor/stem cell has an immunophenotype profile showing positive reactions against antibodies for CD14, CD31, CD44, and-CD45, CD54, CD104, CD105, and CD166 antigens; and negative reactions against antibodies for CD34, CD49a, CD62E, CD73, CD90(Thy-1) and CD133 antigens; positive and partial positive reactions against antibodies for CD54 and CD166 antigens; negative and partial negative reactions against antibodies CD73(SH3, SH4) and CD105(SH2) antigens; and negative and partial positive reactions against antibodies for CD49a and CD104 antigens.

(Canceled)

 (withdrawn – currently amended) An animal cell culture medium composition for isolating and culturing multipotent progenitor/stem cells from cord blood-derived mononuclear cells, which is selected from the group consisting of:

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glutamine; and

an animal cell culture medium composition comprising 10 to 20% FBS, 1 to 2 mM Lglutamine and 10 to 100 ng/M&ml GM-CSF;

an animal cell culture medium composition comprising 10 to 20% FBS and 1 to 2 mM $L\mbox{-}$

an animal cell culture medium composition comprising 10 to 20% FBS, 1 to 2 mM Lglutamine, 10 to 100 ng/mlmg/Mf SCF and 10 to 50 ng/mlmg/Mf EGF,

wherein each of the animal cell culture media further contains inorganic salts, amino acids, and vitamins-and/or-supplementary-factors.

- 8. (withdrawn currently amended): The animal cell culture medium composition of claim 7, wherein the animal cell culture medium further contains D-glucose ranging from 3,500 to 5,500 mg/LMf-and sodium pyruvate ranging from 50 to 200 mg/LMf-
- 9. (Currently amended) A method for differentiating multipotent progenitor/stem cells of claim 5 into neurons, which comprises culturing the multipotent progenitor/stem cells in an animal cell culture medium comprising FBS, L-glutamine, retinoic acid, folskolinfolskolin, nerve growth factor_(NGF), a supplementary element mixture and beta-mercaptocthanol, in addition to D-glucose ranging from 3,500 to 5,500 mg/l mg/Mf-and sodium pyruvate ranging from 50 to 200 mg/l_mg/Mf-.
- 10. (Currently amended) The method of claim 9, wherein the animal cell culture medium contains 0.1 to 2% FBS, 1 to 2 mM L-glutamine, 1 to 25 μ M retinoic acid, 1 to 20 μ M

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folskolin, 10 to 100 ng/mlMt NGF, $4\times$ supplementary element mixture and 1×10^{-6} and 1×10^{-5} % beta-mercaptoethanol.

11. (Currently amended) The method of claim 9, wherein the multipotent progenitor/stem cells are inoculated into the animal cell culture medium at a concentration ranging from 2×10⁴+to 8×10⁴-cells/cm²-1x10⁵ to 4x10⁵ cells/mt and culturing at 37°C under an atmosphere of 5% CO₂ for 1 to 2 weeks.

- 12. (withdrawn currently amended): An animal cell culture medium composition for differentiating multipotent progenitor/stem cells of claim 5 into neurons, which comprises 0.1 to 2% FBS, 1 to 2 mM L-glutamine, 1 to 25 μM retinoic acid, 1 to 20 μM folskolin, 10 to 100 ng/mlM€ NGF, 1× supplementary element mixture and 1×10⁻⁶ to 1×10⁻⁵% beta-mercaptoethanol, in addition to D-glucose ranging from 3,500 to 5,500 mg/LM€ and sodium pyruvate ranging from 50 to 200 mg/L M€.
- 13. (withdrawn currently amended): A method for differentiating multipotent progenitor/stem cells of claim 5 into osteoblasts, which comprises culturing the multipotent progenitor/stem cells in an animal cell culture medium comprising FBS, dexamethason, ascorbate-2-phosphate and β-glycerophosphate, in addition to D-glucose ranging from 3,500 to 5,500 mg/L-M4 and sodium pyruvate ranging from 50 to 200 mg/L-M4.

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14. (withdrawn) The method of claim 13, wherein the animal cell culture medium contains 5 to 20% FBS, 0.1 to 1 μM dexamethasone, 10 to 100 μM ascorbate-2-phosphate and 5 to 20 mM β-glycerophosphate.

- 15. (withdrawn currently amended): The method of claim 13, wherein the multipotent progenitor/stem cells are inoculated into the animal cell culture medium at a concentration ranging from 5×10⁴ to 2×10⁵ cells/em² 2.5x10⁵ to 1x10⁶ cells/ml and culturing at 37°C under an atmosphere of 5% CO₂ for 2 to 3 weeks.
- 16. (withdrawn currently amended): An animal cell culture medium composition for differentiating multipotent progenitor/stem cells of claim 5 into osteoblasts, which comprises 5 to 20% FBS, 0.1 to 1 μ M dexamethasone, 10 to 100 μ M ascorbate-2-phosphate and 5 to 20 mM β -glycerophosphate, in addition to D-glucose ranging from 3,500 to 5,500 mg/LML-and sodium pyruvate ranging from 50 to 200 mg/LML.
- 17. (withdrawn currently amended): A method for differentiating multipotent progenitor/stem cells of claim 5 into endothelial cells, which comprises culturing the multipotent progenitor/stem cells in an animal cell culture medium comprising FBS and vascular endothelial growth factor_(VEGF), in addition to D-glucose ranging from 3,500 to 5,500 mg/LMf- and sodium pyruvate ranging from 50 to 200 mg/LMf-.

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 (withdrawn – currently amended): The method of claim 17, wherein the animal cell culture medium contains 0.1 to 2% FBS and 10 to 100 ng/mlM# VEGF.

- 19. (withdrawn currently amended) The method of claim 17, wherein the multipotent progenitor/stem cells are inoculated into the animal cell culture medium at a concentration ranging from 1×10^6 -to 4×10^6 -cells/em²- 5×10^5 to 2×10^6 cells/mf and culturing at 37° C under an atmosphere of 5% CO₂ for 2 to 3 weeks.
- 20. (withdrawn currently amended) An animal cell culture medium composition for differentiating multipotent progenitor/stem cells of claim 5 into endothelial cells, which comprises 0.1 to 2% FBS and 10 to 100 ng/M&ml VEGF, in addition to D-glucose ranging from 3,500 to 5,500 mg/LM&ml sodium pyruvate ranging from 50 to 200 mg/LM&ml.
- 21. (withdrawn currently amended) A method for differentiating multipotent progenitor/stem cells of claim 5 into myoblasts, which comprises culturing the multipotent progenitor/stem cells in an animal cell culture medium comprising bovine serum albumin_(BSA) and 5-azacytidine, in addition to D-glucose ranging from 3,500 to 5,500 mg/LMf-and sodium pyruvate ranging from 50 to 200 mg/LMf-
- (withdrawn) The method of claim 21, wherein the animal culture medium contains 5 to 10% BSA and 10 to 20 µM 5- azacytidine.

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23. (withdrawn – currently amended) The method of claim 21, wherein the

multipotent progenitor/stem cells are inoculated into the animal cell culture medium at a

concentration ranging from 1×10^5 to 5×10^5 cells/cm² -5×10^5 to 2.5×10^6 cells/mℓ and culturing at

37°C under an atmosphere of 5% CO2 for 5 to 6 weeks.

24. (withdrawn - currently amended) An animal cell culture medium composition

for differentiating multipotent progenitor/stem cells of claim 5 into myoblasts, which comprises

5 to 10% BSA and 10 to 20 μM 5- azacytidine, in addition to D-glucose ranging from 3,500 to

5,500 mg/l Mt-and sodium pyruvate ranging from 50 to 200 mg/l, Mt-

25. (withdrawn – currently amended) A method for differentiating multipotent

progenitor/stem cells of claim 5 into hepatocytes, which comprises culturing the multipotent

progenitor/stem cells in an animal cell culture medium comprising hepatocyte growth factor

(HGF), on costatin M and L-glutamine, in addition to D-glucose ranging from $3{,}500$ to $5{,}500$

mg/<u>l</u> M- ℓ -and sodium pyruvate ranging from 50 to 200 mg/<u>l</u>. M- ℓ -

26. (withdrawn - currently amended) The method of claim 25, wherein the animal

cell culture medium contains 10 to 100 ng/ \underline{ml} Mt HGF, 5 to 50 ng/ \underline{ml} Mt oncostatin M and 1 to 2

mM L-glutamine.

27. (withdrawn – currently amended) The method of claim 25, wherein the

multipotent progenitor/stem cells are inoculated into the animal cell culture medium at a

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concentration ranging from 5×10⁴·to-5×10⁵ eells/em²-2.5x10⁵ to 2.5x10⁶ cells/mf and culturing at 37°C under an atmosphere of 5% CO₂ for 2 to 4 weeks.

28. (withdrawn – currently amended) An animal cell culture medium composition for

differentiating multipotent progenitor/stem cells of claim 5 into hepatocytes, which comprises 10

to 100 ng/mlMt HGF, 5 to 50 ng/mlMt oncostatin M and 1 to 2 mM L-glutamine, in addition to

D-glucose ranging from 3.500 to 5.500 mg/l Mt-and sodium pyruvate ranging from 50 to 200

mg/<u>l.</u>₩€.

29. (withdrawn - currently amended) A method for differentiating multipotent

progenitor/stem cells of claim 5 into dendritic cells, which comprises the steps of: culturing the

multipotent progenitor/stem cells in a first animal cell culture medium comprising FBS, L-

glutamine, GM-CSF and interleukin-4 (IL-4) for inducing immature differentiation; transferring

the immature differentiated cells in a second animal cell culture medium comprising FBS, L-

glutamine, tumor necrosis factor-α (TNF-α), IL-1β, IL-6 and prostaglandin E2; and culturing

them for inducing mature differentiation, wherein each of the animal cell culture media further

contains D-glucose ranging from 3,500 to 5,500 mg/ \underline{L} Mt- and sodium pyruvate ranging from 50

to 200 mg/<u>l.</u> Ml.

30. (withdrawn - currently amended) The method of claim 29, wherein the first

animal cell culture medium contains 1 to 2% FBS, 1 to 2 mM L-glutamine, 10 to 1,000 ng/mlMf

GM-CSF and 10 to 100 ng/ml₩ IL-4, and the second animal cell culture medium contains 1 to

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2% FBS, 1 to 2 mM L-glutamine, 1 to 100 ng/mlM ℓ TNF- α , 1 to 2 ng/mlM ℓ IL-1 β , 100 to 1,000 U/mlM ℓ IL-6 and 0.1 to 10 μ g/mlM ℓ prostaglandin E2.

- 31. (withdrawn currently amended) The method of claim 29, wherein the multipotent progenitor/stem cells are inoculated into the first animal cell culture medium at a concentration ranging from 1×10^5 -to 1×10^7 -cells/cm 3 - 5×10^5 to 5×10^7 cells/ml and culturing at 37°C under an atmosphere of 5% CO₂ for 3 to 15 days, and the immature differentiated cells are culture in the second animal cell culture medium at 37°C under an atmosphere of 5% CO₂ for 1 to 7 days.
- 32. (withdrawn currently amended) An animal cell culture medium composition for differentiating multipotent progenitor/stem cells of claim 5 into dendritic cells, which is selected from the group consisting of:

an animal cell culture medium composition for inducing immature differentiation comprising 1 to 20% FBS, 1 to 2 mM L-glutamine, 10 to 1,000 ng/mlMH GM-CSF and 10 to 100 ng/mlMH IL-4; and

an animal cell culture medium composition for inducing mature differentiation comprising 1 to 20% FBS, 1 to 2 mM L-glutamine, 1 to 100 ng/mlM \pm TNF- α , 1 to 100 ng/mlM \pm IL-1 β , 100 to 10,000 U/mlM \pm IL-6 and 0.1 to 10 μ g/mlM \pm prostaglandin E2,

wherein each of the animal cell culture media further contains D-glucose ranging from 3,500 to 5,500 mg/l MH-and sodium pyruvate ranging from 50 to 200 mg/l, MH-

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33. (Original) A cell composition for a cell therapy comprising the multipotent

progenitor/stem cell of claim 5.

34.

administered into a subject who is in need of treating Parkinson's disease, Alzheimer's diseases, quadriplegia resulting from spinal cord injury, leukemia, apoplexy, encephalophyma, juvenileonset diabetes, cardiac infarction, hepatocirrhosis, muscle diseases, cardiomuscular diseases,

(Currently amended) The cell composition of claim 33, which is used for

liver diseases, blood diseases, $\underline{\text{or the-}} \text{disruption } \underline{\text{and }} \underline{\text{or}} \text{ permanent functional disorder of }$

osteoblasts and or chondrocytes.